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AMENDMENT TO THE CLAIMS

1. (Previously Presented) A medicament for treating addiction craving, characterized in

that the medicament consists of a combination of two administration forms, one of the

administration forms continuously releasing at least one modulator of nicotinic receptors, which

is selected from the group consisting of galanthamine and the pharmacologically acceptable salts

of galanthamine, and the other administration form enabling a rapid entry of galanthamine or one

of its pharmacologically acceptable salts into the central nervous system, wherein the

administration form enabling a quick entry of galanthamine or a pharmacologically acceptable

salt of galanthamine into the central nervous system is selected from the group consisting of:

buccal solutions, spray solutions and drip solutions.

2. (Cancelled).

3. (Previously Presented) The medicament according to claim 1, characterized in that the

administration form continuously releasing the modulator or the modulators of nicotinic

receptors is selected from the group consisting of transdermal therapeutic systems, subcutaneous

implants and intramuscularly injectable preparations.

4. (Previously Presented) The medicament according to claim 3, characterized in that the

intramuscularly injectable preparation is a suspension of microcapsules containing the modulator

or the modulators of nicotinic receptors.

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5. (Previously Presented) The medicament according to claim 3, characterized in that the

administration form continuously releasing the modulator or modulators of nicotinic receptors

releases between 10 mg and 25 mg of galanthamine or a pharmacologically acceptable salt of

galanthamine, per day.

6. (Previously Presented) The medicament according to claim 1, characterized in that the

administration form enabling a quick entry of galanthamine or a pharmacologically acceptable

salt of galanthamine into the central nervous system contains galanthamine or a

pharmacologically acceptable salt of galanthamine in an amount of from 1 to 5 mg.

7. (Cancelled).

8. (Previously Presented) The medicament according to claim 1, characterized in that the

administration form which enables a rapid entry of galanthamine or a pharmacologically

acceptable salt of galanthamine into the central nervous system is in the form of a flexible plastic

container with a capacity of between 1 and 5 ml.

9. (Previously Presented) The medicament according to claim 8, characterized in that the

plastic container is provided with nozzles through which the solution is sprayed or dripped into

the nose.

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10. (Withdrawn) A method for treating substance craving by modulation of neuronal

nicotinic receptors, characterized in that it is a two-stage method wherein a permanent treatment

with a pharmaceutical administration form which continuously delivers a modulator of nicotinic

receptors, which is selected from the group consisting of galanthamine and the

pharmacologically acceptable salts of galanthamine, is supplemented upon the appearance of a

strong craving for a substance by administering galanthamine or a pharmacologically acceptable

salt thereof by means of an administration form which enables rapid entry of galanthamine or of

a pharmaceutically acceptable salt thereof into the central nervous system, wherein the

administration form enabling rapid entry of galanthamine or of a pharmacologically acceptable

salt of galanthamine into the central nervous system is selected from the group consisting of:

buccal solutions, spray solutions and drip solutions.

11. (Withdrawn) The method according to claim 10, characterized in that the substance

craving is a craving for alcoholic beverages and/or tobacco products.

12. (Cancelled).

13. (Withdrawn) The method according to claim 10, characterized in that the

administration form releasing the modulator or the modulators of nicotinic receptors

continuously is selected from the group consisting of transdermal therapeutic systems,

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subcutaneous implants and intramuscularly injectable preparations.

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14. (Withdrawn) The method according to claim 13, characterized in that the

subcutaneously injectable preparation is a suspension of microcapsules containing the modulator

or modulators of nicotinic receptors for intramuscular injection.

15. (Withdrawn) The method according to claim 13, characterized in that the

administration form continuously releasing the modulator or modulators of nicotinic receptors

releases between 10 mg and 25 mg of galanthamine or a pharmacologically acceptable salt of

galanthamine, per day.

16. (Withdrawn) The method according to claim 10, characterized in that the

administration form enabling a quick entry of galanthamine or of a pharmacologically acceptable

salt of galanthamine into the central nervous system contains galanthamine or a

pharmacologically acceptable salt of galanthamine in an amount of from 1 to 5 mg.

17. (Cancelled).

18. (Withdrawn) The method according to claim 10 characterized in that the

administration form which enables a rapid entry of galanthamine or of a pharmacologically

acceptable salt of galanthamine into the central nervous system is in the form of a flexible plastic

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container with a capacity of between 1 and 5 ml.

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- 19. (Withdrawn) The method according to claim 18 characterized in that the plastic container is provided with nozzles through which the solution is sprayed or dripped into the nose.
- 20. (Previously Presented) The medicament according to claim 1, wherein the two administration forms are administered independently.
- 21. (Currently Amended) The medicament according to claim[[2]] 1, wherein the modulator of nicotinic receptors in the administration form continuously releasing the modulator is galanthamine.
- 22. (Withdrawn) The method according to claim 12, wherein the modulator of nicotinic receptors in the administration form continuously releasing the modulator is galanthamine.